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(54) Title: METHODS FOR THE DIAGNOSIS AND TREATMENT OF EPILEPSY

(57) Abstract: A method for the diagnosis of SMEI in a patient comprising detecting an alteration in the SCN1A gene, including in a regulatory region of the gene in a patient sample, and ascertaining whether the alteration is known to be SMEI associated or non-SMEI associated or, if not known to be either, determining the likelihood that it is a SMEI associated alteration.

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METHODS FOR THE DIAGNOSIS AND TREATMENT OF EPILEPSY

Technical Field

The present invention relates to the diagnosis and treatment of epilepsy, particularly severe myoclonic epilepsy of infancy (SMEI).

Background Art

Epilepsies constitute a diverse collection of brain disorders that affect about 3% of the population at some 10 time in their lives (Annegers, 1996). An epileptic seizure can be defined as an episodic change in behaviour caused by the disordered firing of populations of neurons in the central nervous system. This results in varying degrees of involuntary muscle contraction and often a loss of 15 consciousness. Epilepsy syndromes have been classified into more than 40 distinct types based upon characteristic symptoms, types of seizure, cause, age of onset and EEG patterns (Commission on Classification and Terminology of 20 the International League Against Epilepsy, 1989). However the single feature that is common to all syndromes is the persistent increase in neuronal excitability that is both occasionally and unpredictably expressed as a seizure.

A genetic contribution to the aetiology of epilepsy 25 has been estimated to be present in approximately 40% of affected individuals (Gardiner, 2000). As epileptic seizures may be the end-point of a number of molecular aberrations that ultimately disturb neuronal synchrony, genetic basis for epilepsy is likely to heterogeneous. There are over 200 Mendelian diseases which include epilepsy as part of the phenotype. In these diseases, seizures are symptomatic of underlying neurological involvement such as disturbances in brain structure or function. In contrast, there are also a 35 number of "pure" epilepsy syndromes in which epilepsy is the sole manifestation in the affected individuals. These

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are termed idiopathic and account for over 60% of all epilepsy cases.

Idiopathic epilepsies have been further divided into partial and generalized sub-types. Partial (focal local) epileptic fits arise from localized cortical discharges, so that only certain groups of muscles are involved and consciousness may be retained (Sutton, 1990). However, in generalized epilepsy, EEG discharge shows no focus such that all subcortical regions of the brain are involved. Although the observation that generalized epilepsies are frequently inherited is understandable, the mechanism by which genetic defects, presumably expressed constitutively in the brain, give rise to partial seizures is less clear.

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The idiopathic generalized epilepsies (IGE) are the 15 most common group of inherited human epilepsy and do not have simple inheritance. Two broad groups of IGE are now known - the classical idiopathic generalized epilepsies (Commission on Classification and Terminology of the 20 International League Against Epilepsy, 1989) and the newly recognized genetic syndrome of generalized epilepsy with febrile seizures plus (GEFS*) (Scheffer and Berkovic, 1997; Singh et al., 1999).

The classical IGEs are divided into a number of clinically recognizable but overlapping sub-syndromes 25 including childhood absence epilepsy, juvenile absence epilepsy, juvenile myoclonic epilepsy etc (Commission on Classification and Terminology of the International League Against Epilepsy, 1989; Roger et al., 1992). The subsyndromes are identified by age of onset and the pattern of seizure types (absence, myoclonus and tonic-clonic). Some patients, particularly those with tonic-clonic seizures alone do not fit a specifically recognized subsyndrome. Arguments for regarding these as separate 35 syndromes, yet recognizing that they are part of a neurobiological continuum, have been presented previously (Berkovic et al., 1987; 1994; Reutens and Berkovic, 1995).

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GEFS' was originally recognized through large multigeneration families and comprises a variety of subsyndromes. Febrile seizures plus (FS') is a sub-syndrome
where children have febrile seizures occurring outside the
age range of 3 months to 6 years, or have associated
febrile tonic-clonic seizures. Wany family members have a
phenotype indistinguishable from the classical febrile
convulsion syndrome and some have FS' with additional
absence, myoclonic, atonic, or complex partial seizures.
The severe end of the GEFS' spectrum includes myoclonic
astatic epilepsy.

In GEFS* families, linkage analysis on rare multigeneration large families with clinical evidence of a major autosomal dominant gene have demonstrated loci on chromosomes 19q and 2q. Both the 19q and 2q GEFS* loci have been confirmed in independently ascertained large families, and genetic defects have been identified. Families linked to 19q are known and a mutation in the gene for the $\beta 1$ subunit of the neuronal sodium channel (SCN1B) has been identified (Wallace et al., 1998). This mutation results in the loss of a critical disulphide bridge of this regulatory subunit and causes a loss of function in vitro. Families linked to 2q are also known and mutations in the pore-forming α subunit of the neuronal sodium channel (SCN1A) have been identified (FCT/AU01/01648; Escayq et al., 2000).

Severe myoclonic epilepsy of infancy (SMEI) is classed as an epileptic syndrome that manifests as both generalised and focal (partial) seizures (Commission on Classification and Terminology of the International League Against Epilepsy, 1989). SMEI begins with prolonged febrile and afebrile hemiclonic and generalised seizures in the first year of life. Between one and four years, other seizure types evolve including myoclonic, absence and atonic seizures. Neurological development is normal in infancy with progressive slowing after two years. A family history of epilepsy and/or febrile seizures is often found

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in SMEI patients and recent work has shown that family members have epilepsy phenotypes consistent with the GEFS+ spectrum (Singh et al., 2001; Veggiotti, 2001). From a clinical perspective, as GEFS+ and SMEI involve fever-related seizures, it was thought that sodium channel genes may be the target for mutations in SMEI affected individuals. This fact was later confirmed when mutations in the SCN1A gene in SMEI patients were identified (Class et al., 2001; Ohmori et al., 2002). Of interest is that each of these mutations were de novo, a fact difficult to reconcile based on the clinical experience that a significant number of SMEI cases have a family history of GEFS+.

The development of a molecular diagnostic test to aid in the early diagnosis of SMEI is important. Such a test would direct the correct treatment strategy for patients likely to be affected with SMEI and would predict a risk. for seizure aggravation as a result of factors such as fever induced by vaccination or other causes. Clinical studies to determine the molecular basis of SMEI have been variable in their results and have been inconclusive as to single molecular basis for SMEI, particularly as alterations in the SCNIA gene are involved in other epilepsy subtypes. The inventors have recognised the need for such a predictive diagnostic test for SMEI and have 25 therefore established a method that overcomes limitations identified in previous clinical studies and determines the likelihood that an epilepsy patient has SMEI based on a molecular analysis of the SCNIA gene.

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Disclosure of the Invention

In a first aspect of the present invention there is provided a method for the diagnosis of SMEI in a patient comprising detecting an alteration in the SCN1A gene, including in a regulatory region of the gene in a patient sample, and ascertaining whether the alteration is known to be SMEI associated or non-SMEI associated or, if not

known to be either, determining the likelihood that it is a SMEI associated alteration.

This information is important for initiating the correct treatment regimen for a patient. antiepileptic drug (AED) treatments may aggravate seizures in some patients with epilepsy. This may take the form of increased seizure frequency, increased seizure severity, or the appearance of a new seizure type. With respect to SMEI. it is known that carbamazepine, gabapentin, and vigabatrin 10 lamotrigine may aggravate (Bourgeois, 2003) whereas valproate has shown to be of benefit to SMEI patients (Scheffer and Berkovic, 2003). The diagnostic method of the present invention therefore will provide important information towards directing the appropriate primary AED selection in patients suspected of having SMEI.

The nature of the alterations in the SCNIA gene may encompass all forms of gene mutations including deletions, insertions, rearrangements and point mutations in the coding and non-coding regions such as the promoter, introns or untranslated regions. Deletions may be of the entire gene or only a portion of the gene whereas point mutations may result in stop codons, frameshifts or amino acid substitutions. Point mutations occurring in the regulatory regions of SCNIA, such as in the promoter, may lead to loss or a decrease of expression of the mRNA or may abolish proper mRNA processing leading to a decrease in mRNA stability or translation efficiency.

The identification of SCNIA alterations in a patient that lead to more severe changes to the SCNIA protein (such as frameshift mutations and nonsense mutations leading to a truncated protein) increases the likelihood that the patient has SMEI. This likelihood is increased even further if it can be shown that the alteration is a de novo change rather than one that is inherited from the patients parents or relatives, or that the alteration in the SCNIA gene is one that has previously been associated

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with SMEI. The flow chart in Figure 1 illustrates one aspect of the present invention.

In an embodiment there is provided a method for the diagnosis of SMEI in a patient comprising performing one or more assays to test for the existence of an SCMIA alteration and to identify the nature of the alteration.

In a further embodiment there is provided a method for the diagnosis of SMEI in a patient comprising the steps of:

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- performing one or more assays to test for the existence of an alteration in the SCNIA gene of the patient; and, if the results indicate the existence of an alteration in the SCNIA gene,
- (2) performing one or more assays to identify the nature of the SCN1A alteration.

There exists a number of assay systems that can be used to test for the existence of an SCNIA alteration and the invention is not limited by the examples that are provided below.

In one embodiment an assay system employed may be the analysis of SCNIA DNA from a patient sample in comparison to wild-type SCNIA DNA. Genomic DNA may be used for the diagnostic analysis and may be obtained from a number of sources including, but not limited to, body cells, such as those present in the blood or cheek, tissue biopsy, surgical specimen, or autopsy material, The DNA may be isolated and used directly for the diagnostic assays or may be amplified by the polymerase chain reaction (PCR) prior to analysis. Similarly, RNA or cDNA may also be used, with or without PCR amplification. In addition, prenatal diagnosis can be accomplished by testing fetal cells, placental cells or amniotic fluid.

In a specific embodiment, a DNA hybridisation assay may be employed. These may consist of probe-based assays specific for the SCN1A gene. One such assay may look at a series of Southern blots of DNA that has been digested with one or more restriction enzymes. Each blot may

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contain a series of normal individuals and a series of patient samples. Samples displaying hybridisation fragments that differ in length from normal DNA when probed with sequences near or including the SCMLA gene (SCNLA gene probe) indicate a possible SCMLA alteration. If rastriction enzymes that produce very large restriction fragments are used then pulsed field gel electropheresis (PFGE) may be employed.

SCNIA exon specific hybridisation assays may also be employed. This type of probe-based assay will utilize at least one probe which specifically and selectively hybridises to an exon of the SCNIA gene in its wild-type form. Thus, the lack of formation of a duplex nucleic acid hybrid containing the nucleic acid probe is indicative of the presence of an alteration in the SCNIA gene. Because of the high specificity of probe-based tests, any negative result is highly indicative of the presence of an SCNIA alteration however further investigational assays should be employed to identify the nature of the alteration to determine the likelihood it is an SMEI-associated alteration.

The SCN1A exon specific assay approach could also be adapted to identify previously determined alterations responsible for SMEI. In this aspect, a probe which specifically and selectively hybridises with the SCN1A gene in its altered form is used (allele specific probe). In this case the formation of a duplex nucleic acid hybrid containing the nucleic acid probe indicative of the presence of the alteration in the SCN1A gene. In each variation of the exon specific assay approach, it is important to take into account known polymorphisms in the SCN1A gene that are not associated with SMEI. A secondary assay such as DNA sequencing should subsequently be employed to ensure that any suspected alterations are not known polymorphisms.

The SCN1A exon specific probes used for each of the abovementioned assays may be derived from: (1) PCR

amplification of each exon of the SCN1A gene using intron specific primers flanking each exon; (2) cDNA probes specific for each exon; or (3) a series of oligonucleotides that collectively represent an SCN1A exon.

In a further embodiment, an assay to analyse heteroduplex formation may be employed. By mixing denatured wild-type SCNIA DNA with a DNA sample from a patient, any sequence variations in the SCN1A sequence between the two samples will lead to the formation of a mixed population of heteroduplexes and homoduplexes during reannealing of the DNA. Analysis of this mixed population can be achieved through the use of such techniques as high liquid chromatography performance (HPLC) which performed under partially denaturing temperatures. In this manner, heteroduplexes will elute from the HPLC column earlier than the homoduplexes because of their reduced melting temperature.

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In a further embodiment, patient samples may be 20 subject to electrophoretic-based assays. For electrophoretic assays that determine SCN1A fragment length differences may be employed. Fragments of each patient's genomic DNA are amplified with SCN1A gene intron specific primers. The amplified regions of the SCN1A gene 25 therefore include the exon of interest, the splice site junction at the exon/intron boundaries, and a short portion of intron at either end of the amplification product. The amplification products may be run on an electrophoresis size-separation gel and the lengths of the amplified fragments are compared to known and expected 30 standard lengths from the wild-type gene to determine if an insertion or deletion mutation is found in the patient sample. This procedure can advantageously be used in a "multiplexed" format, in which primers for a plurality of 35 exons (generally from 2 to 8) are co-amplified, and evaluated simultaneously on a single electrophoretic gel. This is made possible by careful selection of the primers

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for each exon. The amplified fragments spanning each exon are designed to be of different sizes and therefore distinguishable on an electrophoresis/size separation gel. The use of this technique has the advantage of detecting both normal and mutant alleles in heterozygous individuals. Furthermore, through the use of multiplexing it can be very cost effective.

In a further approach, diagnostic electrophoretic assays for the detection of previously identified SCNIA alterations responsible for SMEI may utilise PCR primers which bind specifically to altered exons of the SCNIA gene. In this case, product will only be observed in the electrophoresis gel if hybridization of the primer occurred. Thus, the appearance of amplification product is an indicator of the presence of the alteration, while the length of the amplification product may indicate the presence of additional alterations.

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Additional electrophoretic assays may be employed. These may include the single-stranded conformational polymorphism (SSCP) procedure (Orita et al., 1989). As 20 mentioned above, fragments of each patient's genomic DNA are PCR amplified with SCN1A gene intron specific primers such that individual exons of the SCNIA gene are amplified and may be analysed individually. Exon-specific PCR products are then subjected to electrophoresis on non-25 denaturing polyacrylamide gels such that DNA fragments migrate through the gel based on their conformation as dictated by their sequence composition. SCN1A exonspecific fragments that vary in sequence from wild-type SCN1A sequence will have a different secondary structure 30 conformation and therefore migrate differently through the gel. Aberrantly migrating PCR products in patient samples are indicative of an alteration in the SCN1A exon and should be analysed further in secondary assays such as DNA sequencing to identify the nature of the alteration. 35

Additional electrophoretic assays that may be employed include RNase protection assays (Finkelstein et

al., 1990; Kinszler et al., 1991) and denaturing gradient gel electrophoresis (DGGE) (Wartell et al., 1990; Sheffield et al., 1989). RNase protection involves cleavage of a mutant polynucleotide into two or more smaller fragments whereas DGGE detects differences in migration rates of mutant sequences compared to wild-type sequences, using a denaturing gradient gel.

In the RNase protection assay a labelled riboprobe which is complementary to the human wild-type SCN1A gene coding sequence is hybridised with either mRNA or DNA 10 isolated from the patient and subsequently digested with the enzyme RNase A which is able to detect some mismatches in a duplex RNA structure. If a mismatch is detected by RNase A, it cleaves at the site of the mismatch. Thus, when the annealed RNA preparation is separated on 15 electrophoretic gel matrix, if a mismatch has been detected and cleaved by RNase A, an RNA product will be seen which is smaller than the full length duplex RNA for the riboprobe and the mRNA or DNA. The riboprobe need not be the full length of the SCN1A mRNA or gene but can be a 20 segment of either. If the riboprobe comprises only a segment of the SCN1A mRNA or gene, it will be desirable to use a number of these probes to screen the whole mRNA sequence for mismatches.

In a further embodiment, enzymatic based assays (Taylor and Deeble, 1999) may be used in diagnostic applications. Such assays include the use of SI nuclease, ribonuclease, T4 endonuclease VII, MutS (Modrich, 1991), Cleavase and MutY. In the MutS assay, the protein binds only to sequences that contain a nucleotide mismatch in a heteroduplex between mutant and wild-type sequences.

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When an assay is to be based upon the SCNIA protein, a variety of approaches are possible. For example, diagnosis can be achieved by monitoring differences in the electrophoretic mobility of normal SCNIA protein and SCNIA protein isolated from a patient sample. Such an approach will be particularly useful in identifying alterations in

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which charge substitutions are present, or in which insertions, deletions or substitutions have resulted in a significant change in the electrophoretic migration of the resultant protein. Alternatively, diagnosis may be based upon differences in the proteclytic cleavage patterns of normal and altered proteins, differences in molar ratios of the various amino acid residues, or by functional assays demonstrating altered function of the gene products.

10 Further assays that are based on the SCN1A protein include immunoassays. Immunoassays for the SCN1A gene product are not currently known. However, immunoassay is included in the selection of assays because the procedures for raising antibodies against specific gene products are 15 well described in the literature, for example in U.S. Pat. Nos. 4,172,124 and 4,474,893 which are incorporated herein by reference. Antibodies are normally raised which bind to portions of the gene product away from common mutation sites such that the same antibody binds to both mutant and 20 normal protein. Preferred antibodies for use in this invention are monoclonal antibodies because of their improved predictability and specificity. It will appreciated, however, that essentially any antibody which possesses the desired high level of specificity can be 25 used, and that optimization to achieve high sensitivity is not required.

For the diagnostic detection of novel alterations in SCN1A involved in SMEI, antibodies raised to the carboxyterminal end of the protein would be preferable. For the diagnostic detection of SCN1A alterations previously identified to be involved in SMEI, antibody raised against the defective gene product is preferable. Antibodies are added to a portion of the patient sample under conditions where an immunological reaction can occur, and the sample is then evaluated to see if such a reaction has occurred. The specific method for carrying out this evaluation is not critical and may include enzyme-linked immunosorbant

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assays (ELISA), described in U.S. Pat. No. 4,016,043, which is incorporated herein by reference; fluorescent enzyme immunoassay (FEIA or ELFA), which is similar to ELISA, except that a fluoregenic enzyme substrate such as 4-methylumbelliferyl-beta-galactoside is used instead of a chromogenic substrate, and radioinmunoassay (RIA).

The most definitive diagnostic assay that may be employed is DNA sequencing, and ultimately may be the only assay that is needed to be performed. Comparison of the SCN1A DNA wild-type sequence with the SCN1A sequence of a 10 test patient provides both high specificity and high sensitivity. The general methodology employed involves amplifying (for example with PCR) the DNA fragments of interest from patient DNA; combining the amplified DNA with a sequencing primer which may be the same as or different from the amplification primers; extending the sequencing primer in the presence of normal nucleotide (A. C, G, and T) and a chain-terminating nucleotide, such as a dideoxynucleotide, which prevents further extension of the primer once incorporated; and analyzing the product for the length of the extended fragments obtained. While such methods, which are based on the original dideoxysequencing method disclosed by Sanger et al., 1977 are useful in the present invention, the final assay is not limited to such methods. For example, other methods for determining the sequence of the gene of interest, or a portion thereof, may also be employed. Alternative methods include those described by Maxam and Gilbert (1977) and variations of the dideoxy method and methods which do not rely on chain-30 terminating nucleotides at all such as that disclosed in U.S. Pat. No. 4,971,903, which is incorporated herein by reference. Any sequence differences (other than benion polymorphisms) in SCN1A exons of a test patient when compared to that of the wild-type SCN1A sequence indicate a potential SMEI-causing alteration. 35

In a further aspect of the invention there is provided a method for the diagnosis of SMEI in a patient

comprising the steps of selecting a system of assays comprising one or more assays to provide a test for the existence of an SCN1A alteration, and one or more assays to provide a test to identify the nature of the alteration, so as to determine the likelihood that it is an SMET-associated alteration.

Application of the invention has lead to the identification of a number of mutations in the SCN1A gene in individuals that have been clinically diagnosed with SMEI. This demonstrates the utility of the diagnostic assay in providing a likelihood that an individual may be affected with SMEI.

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According to a further aspect of the present invention there is provided an isolated nucleic acid molecule encoding an altered SCNIA subunit of a mammalian voltage-gated sodium channel, wherein the alteration gives rise to an SMEI phenotype and has the sequence set forth in one of SEQ ID NOS: 1-25.

In a further aspect of the present invention there is provided an isolated polypeptide, said polypeptide being an altered SCN1A subunit of a mammalian voltage-gated sodium channel, wherein the polypeptide has the amino acid sequence set forth in one of SEQ ID NOS: 26-48 and the alteration gives rise to an SMEI phenotype.

Additional alterations in the SCN1A gene were identified during this study. These alterations were identified in individuals that were not suspected of being affected with SMEI based on a clinical diagnosis.

Accordingly, in a further aspect of the present invention there is provided an isolated nucleic acid molecule encoding an altered SCNLA subunit of a mammalian voltage-gated sodium channel, wherein the alteration gives rise to a non-SMEI epilepsy phenotype and has the sequence set forth in one of SEQ ID NOS: 49-53.

In a still further aspect of the present invention there is provided an isolated polypeptide, said polypeptide being an altered SCNLA subunit of a mammalian

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voltage-gated sodium channel, wherein the polypeptide has the amino acid sequence set forth in one of SEQ ID NOS: 54-58 and the alteration gives rise to a non-SMEI epilepsy phenotype.

5 In another aspect of the present invention there is provided an isolated nucleic acid molecule comprising the nucleotide sequence set forth in any one of SEQ ID NO: 1-25, 49-53.

In another espect of the present invention there is provided an isolated nucleic acid molecule consisting of the nucleotide sequence set forth in any one of SEQ ID NO: 1-25, 49-53.

The nucleotide sequences of the present invention can be engineered using methods accepted in the art for a variety of purposes. These include, but are not limited to, modification of the cloning, processing, and/or expression of the gene product. PCR reassembly of gene fragments and the use of synthetic oligonucleotides allow the engineering of the nucleotide sequences of the present invention. For example, oligonucleotide-mediated site-directed mutagenesis can introduce further mutations that create new restriction sites, alter expression patterns and produce splice variants etc.

As a result of the degeneracy of the genetic code, a number of polynucleotide sequences, some that may have minimal similarity to the polynucleotide sequences of any known and naturally occurring gene, may be produced. Thus, the invention includes each and every possible variation of a polynucleotide sequence that could be made by selecting combinations based on possible codon choices. These combinations are made in accordance with standard triplet genetic code as applied to the polynucleotide sequences of the present invention, and all such variations are to be considered as being specifically disclosed.

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The nucleic acid molecules of this invention are typically DNA molecules, and include cDNA, genomic DNA,

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synthetic forms, and mixed polymers, both sense and antisense strands, and may be chemically or biochemically or may contain non-natural or derivatised nucleotide bases as will be appreciated by those skilled art. Such modifications include methylation. intercalators, alkvlators and modified linkages. In some instances it may be advantageous to produce nucleotide sequences possessing a substantially different codon usage than that of the polynucleotide sequences of the present invention. For example, codons 10 may be selected to increase the rate of expression of the peptide in a particular prokaryotic or eukaryotic host corresponding with the frequency that particular codons are utilized by the host. Other reasons to alter the 15 nucleotide sequence without altering the encoded amino acid sequences include the production of RNA transcripts having more desirable properties, such as a greater halfthan transcripts produced from the naturally occurring mutated sequence.

20 The invention also encompasses production of nucleic. acid sequences of the present invention entirely by synthetic chemistry. Synthetic sequences may be inserted into expression vectors and cell systems that contain the necessary elements for transcriptional and translational 25 control of the inserted coding sequence in a suitable host. These elements may include regulatory sequences. promoters, 5' and 3' untranslated regions and specific initiation signals (such as an ATG initiation codon and Kozak consensus sequence) which allow more efficient 3.0 translation of sequences encoding the polypeptides of the present invention. In cases where the complete coding sequence, including the initiation codon and upstream regulatory sequences, are inserted into the appropriate expression vector, additional control signals may not be needed. However, in cases where only coding sequence, or a fragment thereof, is inserted, exogenous translational control signals as described above should be provided by

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the vector. Such signals may be of various origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of enhancers appropriate for the particular host cell system used (Scharf et al., 1994).

The invention also includes nucleic acid molecules that are the complements of the sequences described herein.

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The present invention allows for the preparation of purified polypeptide or protein from the polynucleotides of the present invention, or variants thereof. In order to do this, host cells may be transformed with a novel nucleic acid molecule as described above. Typically said host cells are transfected with an expression vector comprising a DNA molecule according to the invention. A 15 variety of expression vector/host systems may be utilized to contain and express sequences encoding polypeptides of the invention. These include, but are not limited to, microorganisms such as bacteria transformed with plasmid. or cosmid DNA expression vectors; yeast transformed with veast expression vectors; insect cell systems infected with viral expression vectors (e.g., baculovirus); or mouse or other animal or human tissue cell systems. Mammalian cells can also be used to express a protein using a vaccinia virus expression system. The invention is not limited by the host cell or vector employed.

The polynucleotide sequences, or variants thereof, of the present invention can be stably expressed in cell lines to allow long term production of recombinant proteins in mammalian systems. Sequences encoding the polypeptides of the present invention can be transformed into cell lines using expression vectors which may contain viral origins of replication and/or endogenous expression elements and a selectable marker gene on the same or on a separate vector. The selectable marker confers resistance to a selective agent, and its presence allows growth and recovery of cells which successfully express introduced sequences. Resistant clones of stably

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transformed cells may be propagated using tissue culture techniques appropriate to the cell type.

The protein produced by a transformed cell may be secreted or retained intracellularly depending on the sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing polynucleotides which encode a protein may be designed to contain signal sequences which direct secretion of the protein through a prokaryotic or eukaryotic cell membrane.

In addition, a host cell strain may be chosen for its 10 ability to modulate expression of the inserted sequences or to process the expressed protein in the desired fashion. Such modifications of the polypeptide include, but are not limited to, acetylation, glycosylation, 15 phosphorylation. and acvlation. Post-translational cleavage of a "prepro" form of the protein may also be used to specify protein targeting, folding, and/or activity. Different host cells having specific cellular: machinery and characteristic mechanisms for translational activities (e.g., CHO or HeLa cells), are: 20 available from the American Type Culture Collection (ATCC) and may be chosen to ensure the correct modification and processing of the foreign protein.

When large quantities of the protein product of the gene are needed, such as for antibody production, vectors which direct high levels of expression of this protein may be used, such as those containing the T5 or T7 inducible bacteriophage promoter. The present invention also includes the use of the expression systems described above in generating and isolating fusion proteins which contain important functional domains of the protein. These fusion proteins are used for binding, structural and functional studies as well as for the generation of appropriate antibodies.

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35 In order to express and purify the protein as a fusion protein, the appropriate cDNA sequence is inserted into a vector which contains a nucleotide sequence

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encoding another peptide (for example, glutathionine succinyl transferase). The fusion protein is expressed and recovered from prokaryotic or eukaryotic cells. The fusion protein can then be purified by affinity chromatography based upon the fusion vector sequence. The desired protein is then obtained by enzymatic cleavage of the fusion protein.

Fragments of the polypeptides of the present invention may also be produced by direct peptide synthesis using solid-phase techniques. Automated synthesis may be achieved by using the ABI 431A Peptide Synthesizer (Perkin-Elmer). Various fragments of this protein may be synthesized separately and then combined to produce the full-length molecule.

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15 In another aspect of the present invention there is provided an isolated polypeptide comprising the amino acid sequence set forth in any one of SEQ ID Numbers: 26-48, 54-58.

In another aspect of the present invention there is provided an isolated polypeptide consisting of the amino acid sequence set forth in any one of SEQ ID NO: 26-48, 54-58.

According to still another aspect of the invention, there is provided a mammalian voltage-gated sodium channel that incorporates an altered SCNIA protein as described above.

According to still another aspect of the present invention there is provided an expression vector comprising a nucleic acid molecule as described above.

30 According to still another aspect of the present invention there is provided a cell comprising a nucleic acid molecule as described above.

According to still another aspect of the present invention there is provided a method of preparing a polypeptide, said polypeptide being an altered SCN1A protein of a mammalian voltage-gated sodium channel, comprising the steps of:

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 culturing a cell as described above under conditions effective for polypeptide production;
 and

(2) harvesting the polypeptide.

The mutant SCMIA protein may be allowed to assemble with other subunits of the sodium channel that are co-expressed by the cell (such as the SCMIB protein), whereby the assembled eltered sodium channel is hervested.

According to still another aspect of the invention 10 there is provided a polypeptide which is the product of the process described above.

Substantially purified protein or fragments thereof can then be used in further biochemical analyses to establish secondary and tertiary structure. Such methodology is known in the art and includes, but is not restricted to, X-ray crystallography of crystals of the proteins or of the assembled ion channel incorporating the proteins or by nuclear magnetic resonance (NMR). Determination of structure allows for the rational design of pharmaceuticals to interact with the altered sodium channel as a whole or through interaction with the altered SCN1A protein of the channel (see drug screening below), alter the overall sodium channel protein charge configuration or charge interaction with other proteins, or to alter its function in the cell.

It will be appreciated that having identified novel alterations in the SCMIA gene responsible for epilepsy, including SMEI, the altered SCMIA proteins will enable therapeutic methods for the treatment of epilepsy, including SMEI.

Therapeutic Applications

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According to still another aspect of the invention there is provided a method of treating spilepsy, including SMEI, comprising administering a selective antagonist, agonist or modulator of an SCNIA polypeptide as described above to a subject in need of such treatment. In still another aspect of the invention there is provided the use of a selective antagonist, agonist or modulator of an SCMIA polypeptide as described above in the manufacture of a medicament for the treatment of spilepsy, including SMEI.

In one aspect, a suitable antagonist, agonist or modulator will restore wild-type function to sodium channels containing SCNIA alterations that form part of this invention, or will negate the effects the altered receptor has on cell function.

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Using methods well known in the art, an altered sodium channel, or SCNIA protein of the channel, that is causative of epilepsy, including SMEI, may be used to produce antibodies specific for the altered channel or SCNIA protein of the channel or to screen libraries of pharmaceutical agents to identify those that bind the altered channel or SCNIA protein of the channel.

In one aspect, an antibody, which specifically binds to an altered sodium channel or altered SCNIA protein of withe invention, may be used directly as an agonist, antagonist or modulator, or indirectly as a targeting or delivery mechanism for bringing a pharmaceutical agent to cells or tissues that express the altered channel.

In a still further aspect of the invention there is provided an antibody which is immunologically reactive with a polypeptide as described above, but not with a wild-type SCNIA channel or SCNIA protein thereof.

In particular, there is provided an antibody to an assembled sodium channel containing an alteration in the SCNIA protein that forms part of the channel, which is causative of epilepsy, including SMEI. Such antibodies may include, but are not limited to, polyclonal, monoclonal, chimeric, and single chain antibodies as would be understood by the person skilled in the art.

35 For the production of antibodies, various hosts including rabbits, rats, goats, mice, humans, and others may be immunized by injection with a polypeptide as

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described above or with any fragment or oligopeptide thereof which has immunogenic properties. Various adjuvents may be used to increase immunological response and include, but are not limited to Freund's, mineral gels such as aluminium hydroxide, and surface-active substances such as lysolecithin. Adjuvants used in humans include BCG (bacilli Calmette-Guerin) and Corynebacterium parvum.

It is preferred that the oligopeptides, peptides, or fragments used to induce antibodies to the altered sodium channel, or altered SCNIA protein thereof, have an amino acid sequence consisting of at least 5 amino acids, and, more preferably, of at least 10 amino acids. It is also preferable that these oligopeptides, peptides, or fragments are identical to a portion of the amino acid sequence of the natural protein and contain the entire amino acid sequence of a small, naturally occurring molecule. Short stretches of SCNIA amino acids may be fused with those of another protein, such as KLH, and antibodies to the chimeric molecule may be produced.

Monoclonal antibodies to an altered sodium channel, or altered SCNIA protein thereof, may be prepared using any technique which provides for the production of antibody molecules by continuous cell lines in culture.

These include, but are not limited to, the hybridoma technique, the human B-cell hybridoma technique, and the EBV-hybridoma technique. (For example, see Kohler et al., 1975; Kozbor et al., 1985; Cote et al., 1983; Cole et al., 1984).

30 Monoclonal antibodies produced may include, but are not limited to, mouse-derived antibodies, humanised antibodies and fully human antibodies.

Antibodies may also be produced by inducing in vivo production in the lymphocyte population or by screening is immunoglobulin libraries or panels of highly specific binding reagents as disclosed in the literature. (For

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example, see Orlandi et al., 1989; Winter and Milstein, 1991).

Antibody fragments which contain specific binding sites for an altered sodium channel, or altered SCN1A protein thereof, may also be generated. For example, such fragments include, F(ab')2 fragments produced by papsin digestion of the antibody molecule and Fab fragments generated by reducing the disulfide bridges of the F(ab')2 fragments. Alternatively, Fab expression libraries may be constructed to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity. (For example, see Huse et al., 1989).

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Various immunoassays may be used for screening to identify antibodies having the desired specificity. Numerous protocols for competitive binding either polyclonal immunoradiometric assays using ormonoclonal antibodies with established specificities are: well known in the art. Such immunoassays typically involve: the measurement of complex formation between an ion channel and its specific antibody. A two-site, monoclonal- x based immunoassay utilizing antibodies reactive to two non-interfering sodium channel epitopes is preferred, but a competitive binding assay may also be employed.

In a further aspect of the invention there is provided a method of treating epilepsy, including SMEI comprising administering an isolated nucleic acid molecule which is the complement (antisense) of any one of the nucleic acid molecules described above and which encodes an RNA molecule that hybridizes with the mRNA encoding an 30 altered SCNIA of the invention, to a subject in need of such treatment.

In a still further aspect of the invention there is provided the use of an isolated nucleic acid molecule which is the complement (antisense) of a nucleic acid molecule of the invention and which encodes an RNA molecule that hybridizes with the mRNA encoding an altered

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SCN1A of the invention, in the manufacture of a medicament for the treatment of epilepsy, including SMEI.

Typically, a vector expressing the complement (antisense) of the polynuclsotides of the invention may be administered to a subject in need of such treatment. Many methods for introducing vectors into cells or tissues are available and equally suitable for use in vivo, in vitro, and ex vivo. For ex vivo therapy, vectors may be introduced into stem cells taken from the patient and clonally propagated for autologous transplant back into that same patient. Delivery by transfection, by liposome injections, or by polycationic amino polymers may be achieved using methods which are well known in the art. (For example, see Goldman et al., 1997).

Additional antisense or gene-targeted silencing strategies may include, but are not limited to, the use of antisense oligonucleotides, injection of antisense RNA, transfection of antisense RNA expression vectors, and the use of RNA interference (RNAi) or short interfering RNAs (siRNA). Still further, catalytic nucleic acid molecules, such as DNAzymes and ribozymes may be used for gene silencing (Breaker and Joyce, 1994; Haseloff and Gerlach, 1988). These molecules function by cleaving their target mRNA molecule rather than merely binding to it as in traditional antisense approaches.

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In a further aspect, a suitable agonist, antagonist or modulator may include peptides, phosphopeptides or small organic or inorganic compounds that can restore wild-type activity of sodium channels containing alterations in SCNIA protein of the receptor as described above.

Peptides, phosphopeptides or small organic or inorganic compounds suitable for therapeutic applications may be identified using nucleic acids and peptides of the invention in drug screening applications as described below. Molecules identified from these screens may also be of therapeutic application in affected individuals

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carrying other sodium channel alterations, or individuals carrying alterations in genes other than those comprising the sodium channel, if the molecule is able to correct the common underlying functional deficit imposed by these elterations and those of the invention.

There is therefore provided a method of treating epilepsy, including SMEI comprising administering a compound that is a suitable agonist, antagonist or modulator of a sodium channel and that has been identified using altered SCNIA of the invention.

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In some instances, an appropriate approach for treatment may be combination therapy. This may involve the administering an antibody, an agonist, antagonist or modulator, or complement (antisense) to an altered sodium channel, or altered SCNIA protein thereof, invention to inhibit its functional effect, combined with administration of wild-type SCN1A which may restore levels of wild-type sodium channel formation to normal levels. Wild-type SCN1A can be administered using gene therapy. described above for complement approaches as administration.

There is therefore provided a method of treating epilepsy, including SMEI comprising administration of an antibody, an agonist, antagonist or modulator, or complement to an altered sodium channel, or altered SCNIA protein thereof, of the invention in combination with administration of wild-type SCNIA.

In still another aspect of the invention there is provided the use of an antibody, an agonist, antagonist or modulator, or complement to an altered sodium channel, or altered SCNIA protein thereof, of the invention in combination with the use of wild-type SCNIA, in the manufacture of a medicament for the treatment of epilepsy, including SMEI.

In further embodiments, any of the agonists, antagonists, modulators, antibodies, complementary sequences or vectors of the invention may be administered

alone or in combination with other appropriate therapeutic agents. Selection of the appropriate agents may be made by those skilled in the art, according to conventional pharmsceutical principles. The combination of therapeutic agents may act synergistically to effect the treatment or prevention of the various disorders described above. Using this approach, therapeutic efficacy with lower dosages of each agent may be possible, thus reducing the potential for adverse side effects.

Any of the therapeutic methods described above may be applied to any subject in need of such therapy, including, for example, mammals such as dogs, cats, cows, horses, rabbits, monkeys, and most preferably, humans.

15 Drug Screening

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According to still another aspect of the invention, nucleic acid molecules of the invention as well as peptides of the invention, particularly purified altered SCNIA protein and cells expressing these, are useful for the screening of candidate pharmaceutical compounds for the treatment of epilepsy, including SMBI.

Still further, it provides the use of an altered sodium channel polypeptide complex for the screening of candidate pharmaceutical compounds.

25 Still further, it provides the use wherein high throughput screening techniques are employed.

Compounds that can be screened in accordance with the invention include, but are not limited to peptides (such as soluble peptides), phosphopeptides and small organic or inorganic molecules (such as natural product or synthetic chemical libraries and peptidomimetics).

In one embodiment, a screening assay may include a cell-based assay utilising eukaryotic or prokaryotic host cells that are stably transformed with recombinant molecules expressing the polypeptides or fragments of the invention, in competitive binding assays. Binding assays will measure the formation of complexes between an altered

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sodium channel, or altered SCN1A protein thereof, and the compound being tested, or will measure the degree to which a compound being tested will inhibit or restore the formation of a complex between an altered sodium channel, or altered SCM1A protein thereof, and its interactor or licend.

The invention is particularly useful for screening compounds by using the polypeptides of the invention in transformed cells, transfected or injected occytes, or 10 animal models bearing altered SCN1A such as transgenic animals or gene targeted (knock-in) animals transformed hosts). Drug candidates can be added cultured cells that express an altered SCNIA protein (appropriate wild-type sodium channel subunits such as SCN1B should also be expressed for receptor assembly), can 15 be added to occytes transfected or injected with altered SCN1A protein (appropriate wild-type sodium channel subunits such as SCN1B must also be injected for receptor assembly), or can be administered to an animal 20 model expressing an altered SCN1A protein. Determining the ability of the test compound to modulate altered sodium channel activity can be accomplished by a number of techniques known in the art. These include for example measuring the effect on the current of the channel as compared to the current of a cell or animal containing the 25 wild-type sodium channel.

Current in cells can be measured by a number of approaches including the patch-clamp technique (methods described in Hamill et al, 1981) or using fluorescence based assays as are known in the art (see Gonzalez et al., 1999). Drug candidates that alter the current to a more normal level are useful for treating or preventing epilepsy, including SMEI.

Non cell-based assays may also be used for identifying compounds that can inhibit or restore binding between the altered sodium channel, or altered SCNIA protein thereof, of the invention, and their interactors.

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Such assays are known in the art and include for example AlphaScreen technology (PerkinElmer Life Sciences, USA). This application relies on the use of beads such that each interaction partner is bound to a separate bead via an antibody. Interaction of each partner will bring the beads into proximity, such that laser excitation initiates a number of chemical reactions ultimately leading to fluorophores emitting a light signal. Candidate compounds that inhibit the binding of the altered sodium channel, or altered SCN1A protein thereof, with its 1.0 interactor will result in loss of light emission, while candidate compounds that restore the binding of the altered sodium channel, or altered SCNIA protein thereof, with its interactor will result in positive light emission. These assays ultimately enable identification 15 and isolation of the candidate compounds.

High-throughput drug screening techniques may also employ methods as described in W084/03564. Small peptide test compounds synthesised on a solid substrate can be assayed for altered SCNIA protein or altered sodium channel binding. Bound altered sodium channel or altered solim in the art. In a variation of this technique, purified polypeptides of the invention can be coated directly onto plates to identify interacting test compounds.

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The invention also contemplates the use of competition drug screening assays in which neutralizing antibodies capable of specifically binding the altered sodium channel compete with a test compound for binding thereto. In this manner, the antibodies can be used to detect the presence of any peptide that shares one or more antigenic determinants of the altered receptor.

The polypeptides of the present invention may also be used for screening compounds developed as a result of combinatorial library technology. This provides a way to test a large number of different substances for their ability to modulate activity of a polypeptide. A substance

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identified as a modulator of polypeptide function may be peptide or non-peptide in nature. Non-peptide "small molecules" are often preferred for many in vivo pharmaceutical applications. In addition, a mimic be designed οf the substance may pharmaceutical use. The design of mimetics based on a known pharmaceutically active compound ("lead" compound) common approach to the development of novel pharmaceuticals. This is often desirable where the original active compound is difficult or expensive to 10 synthesise or where it provides an unsuitable method of administration. In the design of a mimetic, particular parts of the original active compound that are important in determining the target property are identified. These 15 parts or residues constituting the active region of the compound are known as its pharmacophore. Once found, the pharmacophore structure is modelled according to its physical properties using data from a range of sources including x-ray diffraction data and NMR. A template 20 molecule is then selected onto which chemical groups which mimic the pharmacophore can be added. The selection can be made such that the mimetic is easy to synthesise, is likely to be pharmacologically acceptable, does not degrade in vivo and retains the biological activity of the 25 lead compound. Further optimisation or modification can be carried out to select one or more final mimetics useful for in vivo or clinical testing.

It is also possible to isolate a target-specific antibody and then solve its crystal structure. principle, this approach yields a pharmacophore upon which subsequent drug design can be based as described above. It possible to avoid protein crystallography altogether by generating anti-idiotypic antibodies (antiids) to a functional, pharmacologically active antibody. 35 As a mirror image of a mirror image, the binding site of the anti-ids would be expected to be an analogue of the original receptor. The anti-id could then be used to

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isolate peptides from chemically or biologically produced peptide banks.

Another alternative method for drug screening relies on structure-based rational drug design. Determination of the three dimensional structure of the polypeptides of the invention, or the three dimensional structure of the GABA-B receptors which incorporate these polypeptides allows for structure-based drug design to identify biologically active lead compounds.

Three dimensional structural models can be generated 10 by a number of applications, some of which include experimental models such as x-ray crystallography and NMR and/or from in silico studies of structural databases such the Protein Databank (PDB). In addition, three 15 dimensional structural models can be determined using a number of known protein structure prediction techniques based on the primary sequences of the polypeptides (e.g. SYBYL - Tripos Associated, St. Louis, MO), de novo protein structure design programs (e.g. MODELER - MSI Inc., San. Diego, CA, or MOE - Chemical Computing Group, Montreal, 20 Canada) or ab initio methods as described, for example, in US Patent Numbers 5331573 and 5579250, the contents of which are incorporated herein by reference.

Once the three dimensional structure of a polypeptide or polypeptide complex has been determined, structurebased drug discovery techniques can be employed to design biologically-active compounds based on these three dimensional structures. Such techniques are known in the art and include examples such as DOCK (University of 30 California, San Francisco) or AUTODOCK (Scripps Research Institute, La Jolla, California). A computational docking protocol will identify the active site or sites that are deemed important for protein activity based on a predicted protein model. Molecular databases, such as the Available Chemicals Directory (ACD) are then screened for molecules that complement the protein model.

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Using methods such as these, potential clinical drug candidates can be identified and computationally ranked in order to reduce the time and expense associated with typical 'wet lab' drug screening methodologies.

Compounds identified through screening procedures as described above, and which are based on the use of the altered nucleic acid and polypeptides of the invention, can also be tested for their effect on correcting the functional deficit imposed by other gene alterations in affected individuals including other SCN1A alterations. 10

Such compounds form a part of the present invention, as do pharmaceutical compositions containing these and a pharmaceutically acceptable carrier.

15 Pharmaceutical Preparations

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Compounds identified from screening assays and shown to restore sodium channel wild-type activity can be administered to a patient at a therapeutically effective dose to treat or ameliorate epilepsy, including SMEI, as described above. A therapeutically effective dose refers to that amount of the compound sufficient to result in amelioration of symptoms of the disorder.

Toxicity and therapeutic efficacy of such compounds can be determined by standard pharmaceutical procedures in cell cultures or experimental animals. The data obtained from these studies can then be used in the formulation of a range of dosages for use in humans.

Pharmaceutical compositions for use in accordance with the present invention can be formulated in a conventional manner using one or more physiological acceptable carriers, excipients or stabilisers which are well known. Acceptable carriers, excipients or stabilizers are non-toxic at the dosages and concentrations employed, and include buffers such as phosphate, citrate, and other 35 organic acids; antioxidants including absorbic acid; low molecular weight (less than about 10 polypeptides; proteins, such as serum albumin, gelatin, or

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immunoglobulins; binding agents including hydrophilic polymers such as polyvinylpyrrolidone; amino acids such as glycine, glutamine, asparagine, arginine or lysine; monosaccharides, disaccharides, and other carbohydrates including glucose, mannose, or dextrins; chelating agents such as EDTA; sugar alcohols such as mannitol or sorbitol; salt-forming counterions such as sodium; and/or non-ionic surfactants such as Tween, Pluronics or polyethylene clycol (PEG).

The formulation of pharmaceutical compositions for use in accordance with the present invention will be based on the proposed route of administration. Routes of administration may include, but are not limited to, inhalation, insufflation (either through the mouth or nose), oral, buccal, rectal or parental administration.

Microarray

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In further embodiments, complete cDNAs, coligonucleotides or longer fragments derived from any of the SCN1A polynucleotide sequences described herein may be used as probes in a microarray. The microarray can be used to diagnose epilepsy, including SMEI, through the identification of the SCN1A alterations of the invention, to understand the genetic basis of epilepsy, or can be used to develop and monitor the activities of therapeutic agents.

According to a further aspect of the present invention, tissue material obtained from animal models (see below) generated as a result of the identification of specific SCNIA human alterations of the present invention, can be used in microarray experiments. These experiments can be conducted to identify the level of expression of SCNIA, or the level of expression of any cDNA clone from whole-tissue libraries, in diseased tissue as opposed to normal control tissue. Variations in the expression level of genes, including SCNIA, between the two tissues indicates their possible involvement in the disease

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process either as a cause or consequence of the original SCNIA alteration present in the animal model. These experiments may also be used to determine gene function, to understand the genetic basis of epilepsy, to diagnose spilepsy, and to develop and monitor the activities of therapeutic agents. Microarrays may be prepared, used, and analyzed using methods known in the art. (For example, see Schena et al., 1996; Heller et al., 1997).

10 Transformed Hosts

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The present invention also provides for genetically modified (knock-out, knock-in and transgenic), non-human animal models comprising nucleic acid molecules of the invention. These animals are useful for the study of the function of a sodium channel, to study the mechanisms of epilepsy as related to a sodium channel, for the screening of candidate pharmaceutical compounds, for the creation of explanted mammalian cell cultures which express altered sodium channels, and for the evaluation of potential, therapeutic interventions.

Animal species which are suitable for use in the animal models of the present invention include, but are not limited to, rats, mice, hamsters, guinea pigs, rabbits, dogs, cats, goats, sheep, pigs, and non-human primates such as monkeys and chimpanzees. For initial studies, genetically modified mice and rats are highly desirable due to the relative ease in generating knock-in, knock-out or transgenics of these animals, their ease of maintenance and their shorter life spans. For certain studies, transgenic yeast or invertebrates may be suitable and preferred because they allow for rapid screening and provide for much easier handling. For longer term studies, non-human primates may be desired due to their similarity with humans.

To create an animal model for an altered sodium channel of the invention, several methods can be employed.

These include, but are not limited to, generation of a

specific alteration in a homologous animal gene, insertion of a wild type human gene and/or a humanized animal gene by homologous recombination, insertion of an altered human gene as genomic or minigene cDMA constructs using wild type or altered or artificial promoter elements, insertion of artificially modified fragments of the. homologous recombination. endogenous gene bv The modifications include insertion of mutant stop codons, the inclusion DNA sequences, or the recombination elements (lox p sites) recognized by enzymes such as Cre recombinase.

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To create transgenic mice in order to study gain of gene function in vivo, a SCN1A alteration of the invention can be inserted into a mouse germ line using standard techniques such as cocyte microinjection. Gain of gene function can mean the over-expression of a gene and its protein product, or the genetic complementation of a mutation of the gene under investigation. For cocyte injection, one or more copies of the mutant gene can be inserted into the pronucleus of a just-fertilized mouse $\boldsymbol{\theta}_{\boldsymbol{\theta}}$ 20 occyte. This occyte is then reimplanted into a pseudopregnant foster mother. The live-born mice can then be screened for integrants using analysis of tail DNA for the presence of the relevant human SCNIA gene sequence. The transgene can be either a complete genomic sequence injected as a YAC, BAC, PAC or other chromosome DNA fragment, a cDNA with either the natural promoter or a heterologous promoter, or a minigene containing all of the coding region and other elements found to be necessary for 30 optimum expression.

To generate knock-out mice or knock-in mice, gene targeting through homologous recombination in mouse embryonic stem (ES) cells may be applied. Knock-out mice are generated to study loss of gene function in vivo while knock-in mice (which are preferred) allow the study of gain of function or to study the effect of specific gene mutations. Knock-in mice are similar to transgenic mice

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however the integration site and copy number are defined in the former.

For knock-out mouse generation, gene targeting vectors can be designed such that they delete (knock-out) the protein coding sequence of the SCMIA gene in the mouse genome. In contrast, knock-in mice can be produced whereby a gene targeting vector containing the relevant altered SCMIA gene can integrate into a defined genetic locus in the mouse genome. For both applications, homologous recombination is catalysed by specific DNA repair enzymes that recognise homologous DNA sequences and exchange them via double crossover.

Gene targeting vectors are usually introduced into ES cells using electroporation. ES cell integrants are then isolated via an antibiotic resistance gene present on the targeting vector and are subsequently genotyped to identify those ES cell clones in which the gene under investigation has integrated into the locus of interest. The appropriate ES cells are then transmitted through the germline to produce a novel mouse strain.

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In instances where gene ablation results in early embryonic lethality, conditional gene targeting may be employed. This allows genes to be deleted in a temporally and spatially controlled fashion. As above, appropriate ES cells are transmitted through the germline to produce a novel mouse strain, however the actual deletion of the gene is performed in the adult mouse in a tissue specific or time controlled manner. Conditional gene targeting is most commonly achieved by use of the cre/lox system. The enzyme cre is able to recognise the 34 base pair loxP sequence such that loxP flanked (or floxed) DNA is recognised and excised by cre. Tissue specific cre expression in transgenic mice enables the generation of tissue specific knock-out mice by mating gene targeted floxed mice with cre transgenic mice. Knock-out can be conducted in every tissue (Schwenk et al., 1995) using the 'deleter' mouse or using transgenic mice with an inducible

cre gene (such as those with tetracycline inducible cre genes), or knock-out can be tissue specific for example through the use of the CD19-cre mouse (Rickert et al., 1997).

According to still another aspect of the invention 5 there is provided the use of genetically modified nonhuman animals as described above for the screening of candidate pharmaceutical compounds (see drug screening above). These animals are also useful for the evaluation efficacy, toxicity, metabolism) therapeutic 10 including those candidate pharmaceutical compounds, identified from the invention as described above, for the treatment of epilepsy, including SMEI.

Throughout this specification and the claims, the words "comprise", "comprises" and "comprising" are used in a non-exclusive sense, except where the context requires otherwise.

It will be apparent to the person skilled in the art that while the invention has been described in some detail for the purposes of clarity and understanding, various modifications and alterations to the embodiments and methods described herein may be made without departing from the scope of the inventive concept disclosed in this specification.

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Modes for Performing the Invention

Any combination of assay systems described above may be employed for the identification of SCNIA mutations potentially causative of SMEI. Provided below are examples of assays that may be employed.

Example 1: Patient DNA collection

The flowchart in Figure 1 illustrates a strategy based on the invention that can be used to determine the likelihood that an alteration in the SCNIA gene is responsible for SMEI. The assay combination chosen is preceded by selecting the patient population to be

examined and obtaining DNA from the sample population. The sample population may encompass any individual with epilepsy but would likely focus on children with febrile seizures as well as other patients that are suspected to have myoclonic epilepsy. For the present study, the patient population chosen included individuals that had been diagnosed with SMEI from a clinical analysis or had severe encephalopathies occurring during the first 12 months of life.

DNA from a test patient may be obtained in a number 10 of ways. The most common approach is to obtain DNA from blood samples taken from the patient, however DNA may also be obtained using less invasive approaches such as from cheek cell swabs.

For the current study DNA was extracted from collected blood using the QIAamp DNA Blood Maxi kit (Olagen) according to manufacturers specifications or through procedures adapted from Wyman and White (1980). For DNA samples obtained using the QIAamp kit, a final ethanol precipitation step was employed with DNA pellets; being resuspended in sterile water. Stock DNA samples were kept at a concentration of 200 ng/ul and 100 ng/ul dilutions were prepared for subsequent PCR reactions.

Example 2: dHPLC Assav 25

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Once DNA was obtained from the patients, amplification of individual exons of the SCN1A gene was employed prior to analysis by high performance liquid chromatography (dHPLC). The SCNIA gene has 26 exons for 30 which primers were designed to amplify 33 amplicons. Each exon was amplified by a single amplicon except for exons 11, 15 and 16 which are amplified in two amplicons respectively and exon 26 where 5 amplicons were used to amplify the entire exon. Table 1 provides a list of primers that were designed to analyse each exon of the SCN1A gene.

PCR amplification reactions were performed in a

volume of 20 ul and were prepared in 96-well plates. For the majority of amplicons the PCR reaction consisted of 1X PCR buffer (Invitrogen), 200 uM dNTPs, 300 ng of each primer, 1.5 mM MgCl₂, 100 ng DNA and 0.5 units of Taq DNA polymerase (Invitrogen). The above conditions were used for all amplicons except for except 5, and 26(1) where 1 Unit of Taq DNA polymerase was used.

The thermal cycling conditions employed for PCR amplification varied according to each exon. For exons 1-4, 6-9, 11(1), 11(2), 12, 14, 15(1), 15(2), 16(2), 19, and 22-24, PCR reactions were performed using 1 cycle of 94°C for 2 minutes, followed by 10 cycles of 60°C for 30 seconds, 72°C for 30 seconds, and 94°C for 30 seconds, 72°C for 30 seconds, and 94°C for 30 seconds. A final annealing reaction at 55°C for 30 seconds followed by an extension reaction for 10 minutes at 72°C completed the cycling conditions for these amplicons.

For exon 5, the same conditions were employed as above except the annealing temperature was 62°C for 10, cycles and then 58°C for 25 cycles.

For exons 10, 16(1), 21, 25, 26(1), 26(2), 26(3), 26(4), and 26(5), PCR reactions were performed using 1 cycle of 94°C for 2 minutes, followed by 10 cycles of 60°C for 1.5 minutes, 72°C for 1.5 minutes, and 94°C for 1.5 minutes, followed by 25 cycles of 55°C for 1.5 minutes, 72°C for 1.5 minutes, and 94°C for 1.5 minutes. A final annealing reaction at 55°C for 1.5 minutes followed by an extension reaction for 10 minutes at 72°C completed the cycling conditions for these amplicons.

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For exons 17, 18 and 20, PCR reactions were performed using 1 cycle of 94°C for 2 minutes, followed by 35 cycles of 50°C for 30 seconds, 72°C for 30 seconds, and 94°C for 30 seconds. A final annealing reaction at 50°C for 30 seconds followed by an extension reaction for 10 minutes at 72°C completed the cycling conditions for these amplicons.

For exon 13, PCR reactions were performed using 1 cycle of 94°C for 2 minutes, followed by 10 cycles of 94°C for 1 minute, 64°C for 1.5 minutes, and 72°C for 1.5 minutes, followed by 25 cycles of 94°C for 1 minute, 60°C for 1.5 minutes, and 72°C for 1.5 minutes. This was followed by a final extension reaction for 10 minutes at 72°C to complete the cycling conditions for this amplicon.

Prior to dHPLC analysis, PCR products were heated to 95°C for 5 minutes and are then slowly cooled at -3°C increments for 1.5 minutes (until 25°C is reached). This is to allow the formation of hetero- and homoduplexes depending upon the nucleotide constitution of the PCR product.

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Various dHPLC systems can be used for heteroduplex analysis and mutation detection. This study used the Transgenomic WAVE® System and the methodology supplied with the system. In order to detect mutations on the dHPLC each product needed to be run under partially denaturing conditions. Due to each amplicon of the SCNIA gene having a different sequence, the temperature(s) at which each product is partially denatured needed to be calculated. Using the Transgenomic software supplied with the dHPLC system the required temperatures for each of the amplicons was determined and is shown in Table 2.

Amplicons are fed through the dHPLC column according to manufacturers conditions and computer generated chromatograms are compared between patient samples and wild-type samples. The analysis is done by visually looking at the chromatograms and also using the mutation detection Transgenomic software supplied with the HPLC. Those patient samples showing different peak patterns to wild-type are considered to contain alterations in the SCNIA amplicon under investigation and the DNA from those individuals was subject to a further assay, namely DNA sequencing (see example 3 below), to determine the nature of the SCNIA alteration and to predict the likelihood that the alteration was responsible for SMEI.

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Example 3: DNA Sequencing Assay

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PCR products from the dHPLC analysis that showed different peak patterns to wild-type may be subject to secondary assays such as DNA sequencing to identify the nature of the alteration. In the present study DNA involved This first sequencing was employed. amplification of the amplicon displaying an altered dHPLC chromatogram from the relevant individual followed by purification of the PCR amplified templates for sequencing using QiaQuick PCR preps (Qiagen) based on manufacturers 10 procedures. The primers used to sequence the purified amplicons were identical to those used for the initial amplification step. For each sequencing reaction, 25 ng of primer and 100 ng of purified PCR template were used. The BigDye sequencing kit (ABI) was used for all sequencing 15 reactions according to the manufacturers specifications. The products were run on an ABI 377 Sequencer and analysed. using the EditView program.

A comparison of the DNA sequence obtained from the patient sample was then made directly to that of the wild-type SCNIA sequence in order to identify the nature of the DNA alteration that lead to the change detected by dHPLC.

The results of the screening of 26 of the 33 amplicons of the SCNIA gene are shown in Table 3. A total of 96 patients were analysed with their clinical epilepsy phenotype being hidden during the analysis. A total of 34 samples were shown to have an alteration in the SCNIA gene and of these, 28 samples had a clear SMEI phenotype based on a clinical analysis. Four of the SCNIA alterations (M1780T, R222X, R1407X, R1892X) that were identified are not shown in Table 3 as they had previously been associated with SMEI (Nabbout et al., 2003; Claes et al., 2001; Sugawara et al., 2002). It can therefore be determined that if an SCNIA alteration is found in a patient, then the patient has an 82% chance (28/34) of having SMEI.

This likelihood would increase if the alteration

identified was one that had previously been associated with SMEI. In addition, based on current opinion (Mulley et al., 2003) the likelihood would further increase if the alteration is not seen in the parents or relatives of the affected individual (i.e. is a de novo alteration) and is still further increased if the alteration is found to result in a major disruption to the protein (such as a truncating alteration). The ability to provide this level of certainty as to a diagnosis of SMEI will be of benefit when considering therapy regimes for the patient and the avoidance of seizure aggravation induced by such factors as fever associated with vaccinations and other causes.

Example 4: Additional Assays - SSCP Assay

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In addition to the assays described above, other assays may be employed to test for the existence of alterations in the SCNIA gene that are associated with SMET. One such assay is single strand conformation polymorphism (SSCP) analysis. In this technique, DNA obtained from the patient is first PCR amplified for individual exons of the SCNIA gene. The primers employed for dHPLC analysis (see Table 1) may also be used for SSCP analysis.

In some instances the primers used for SSCP analysis are labelled at their 5' end with HEX for a fluorescent-based detection approach as used for example in the Gelscan 2000 system (Corbett Research, Australia). SSCP PCR reactions and cycling conditions can be performed as described above for dHPLC analysis, however any PCR reaction and cycling conditions may be employed provided that the amplification produces a distinct product specific for the amplicon under investigation only.

An example of alternative PCR reaction conditions are where the reaction is performed in a total volume of 10 μ l containing 67 mM Tris-HCl (pH 8.8); 16.5 mM (NH₄)₂SO₄; 6.5 μ M EDTA; 1.5 mM MgCl₂; 200 μ M each dNTP; 10% DMSO; 0.17 mg/ml BSA; 10 mM β -mercaptoethanol; 5 μ g/ml each primer and

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100 U/ml Taq DNA polymerase. PCR cycling conditions may use 10 cycles of 94°C for 30 seconds, 60°C for 30 seconds, and 72°C for 30 seconds followed by 25 cycles of 94°C for 30 seconds, 55°C for 30 seconds, and 72°C for 30 seconds. A final extension reaction for 10 minutes at 72°C should follow.

Twenty ul of loading dve comprising 50% (v/v) formamide, 12.5 mM EDTA and 0.02% (w/v) bromophenol blue then added to completed reactions which subsequently run on non-denaturing 4% polyacrylamide gels 10 with a cross-linking ratio of 35:1 (acrylamide:bisacrylamide) and containing 2% glycerol. For analysis of PCR amplicons using the GelScan 2000 system, the gel thickness typically employed is 100µm, with a width of 168mm and length of 160mm. Gels are normally run at 1200 volts and approximately 20mA, at 22°C and analysed on the GelScan 2000 system according to manufacturers specifications. Those amplicons that contain alterations in the SCN1A sequence will migrate through the gel differently than wild-type amplicons due to their altered single strand conformation. A further assay such as DNA sequencing may then be employed (see example 3 above) to determine the nature of the SCN1A alteration in the amplicon.

TABLE 1

Primer Sequences Used for dHPLC Assay Analysis of SCN1A

Exon	Forward Primer	Reverse Primer	Size(bp)
1	CCTCTAGCTCATGTTTCATGAC	TGCAGTAGGCAATTAGCAGC	448
2	CTAATTAAGAAGAGATCCAGTGACAG	GCTATAAAGTGCTTACAGATCATGTAC	356
3	CCCTGAATTTTGGCTAAGCTGCAG	CTACATTAAGACACAGTTTCAAAATCC	263
4	GGGCTACGTTTCATTTGTATG	GCAACCTATTCTTAAAGCATAAGACTG	358
5	AGGCTCTTTGTACCTACAGC	CATGTAGGGTCCGTCTCATT	200
6	CACACGTGTTAAGTCTTCATAGT	AGCCCCTCAAGTATTTATCCT	394
7	GAACCTGACCTTCCTGTTCTC	GTTGGCTGTTATCTTCAGTTTC	241
8	AAAGGCAGCAGAACGACTTG	GGATAGAGGAACTCAAGTCTC	322
9	TTGAAAGTTGAAGCCACCAC	CCACCTGCTCTTAGGTACTC	363
10	GCCATGCAAATACTTCAGCCC	CACAACAGTGGTTGATTCAGTTG	480
11(1)	TGAATGCTGAAATCTCCTTCTAC	CTCAGGTTGCTGTTGCGTCTC	306
11(2)	GATAACGAGAGCCGTAGAGAT	TCTGTAGAAACACTGGCTGG	315
12	CATGAAATTCACTGTGTCACC	CAGCTCTTGAATTAGACTGTC	347
13	ATCCTTGGGAGGTTTAGAGT	GCATGAAGGATGGTTGAAAG	510
14	CATTGTGGGAAAATAGCATAAGC	GCTATGCAGAACCCTGATTG	339
15(1)	TGAGACGGTTAGGGCAGATC	AGAAGTCATTCATGTGCCAGC	348
15(2)	GTCTTGGCCATCATCGTCTTC	ACATGTGCACAATGTGCAGG	350
16(1)	GTGGTGTTTCCTTCTCATCAAG	CACTGCTGCCAGTTCCTATAC	458
16(2)	CAACAGTCCTTCATTAGGAAAC	ACCTTCCCACACCTATAGAATC	353
17	CTTGGCAGGCAACTTATTACC	CAAGCTGCACTCCAAATGAAAG	232
18	TGGAAGCAGAGACACTTTATCTAC	GTGCTGTATCACCTTTTCTTAATC	234
19	CCTATTCCAATGAAATGTCATATG	CAAGCTACCTTGAACAGAGAC	318 .
20	CTACACATTGAATGATGATTCTGT	GCTATATACAATACTTCAGGTTCT	216
21	ACCAGAGATTACTAGGGGAAT	CTGGGCTCATAAACTTGTACTAAC	513
22	ACTGTCTTGGTCCAAAATCTG	TTCGATTAATTTTACCACCTGATC	267
23	AGCACCAGTGACATTTCCAAC	GGCAGAGAAAACACTCCAAGG	271
24	GACACAGTTTTAACCAGTTTG	TGTGAGACAAGCATGCAAGTT	207
25	CAGGGCCAATGACTACTTTGC	CTGATTGCTGGGATGATCTTGAATC	477
26(1)	CAGGACTCTGAACCTTACCTTG	ATTCCAACAGATGGGTTCCCA	534
26(2)	TCCTGCGTTGTTTAACATCGG	AGCGCAGCTGCAAACTGAGAT	504
26(3)	TGGAAGCTCAGTTAAGGGAGA	GTAGTGATTGGCTGATAGGAG	480
26(4)	CCGATGCAACTCAGTTCATGGA	TGCCTTCTTGCTCATGTTTTTCCACA	555
26(5)	AGAGCGATTCATGGCTTCCAATCC	TGCTGACAAGGGGTCACTGTCT	526

26(5) AGAGCGATTCATGGCTTCCAATCC TGCTGACAAGGGGTCACTGTCT 526

Note: Primer sequences are listed 5° to 3°. Due to the large size of exons 11, 15, 16, and 26, the exons were split into two or more overlapping amplicons.

TABLE 2

Partial Denaturing Conditions for dHPLC Assay Analysis of SCN1A Amplicons

Exon	Temp 1	Temp 2	Temp 3	Exon	Temp 1	Temp 2	Temp 3
1	53.0	55.2	58.8	16(1)	54.9	55.6	57.3
2	53.5	55.2	58.0	16(2)	56.1	-	-
3	55.9	-	-	17	57.6	60.4	-
4	54.5	55.5	56.5	18	58.5	-	-
5	60.2	-	-	19	53.0	56.5	-
6	53.8	57.5	58.9	20	58.5	-	-
7	56.3	-	-	21	55.2	56.7	-
8	57.9	_	-	22	55.5	-	-
9	56.7	60.1	-	23	55.6	56.3	-
10	56.0	58.5	61.0	24	55.6	56.7	-
11(1)	57.1	60.2	61.5	25	53.8	55.6	56.8
11(2)	58.8	61.2	62.3	26(1)	55.8	59.0	60.0
12	55.3	57.3	-	26(2)	58.5	-	-
13	53.8	55.2	56.4	26(3)	58.5	59.8	-
14	55.4	57.9	-	26(4)	55.5	57.0	57.9
15(1)	57.5	60.2	-	26(5)	55.1	56.6	-
15(2)	58.4	60.7	_				

15(2) 58.4 60.7
Note: All temperatures are in degrees celcius. Temp 1, Temp 2, and Temp 3 represent the temperatures at which different regions of the amplicon denature during the dHPLC analysis. Some amplicons required 3 partially denaturing temperatures for complete analysis of the amplicon whereas other amplicons required two or less temperatures.

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TABLE 3

Novel alterations identified in SCN1A Nucleotide Amino Acid SEO ID Patient Mutation Diagnosis1 Change² Change² Type Numbers SMEI Missense Y84C 1, 26 c251A→G SMEI Missense c301C→T R101W 2, 27 SMEI Missense 1171K 3, 28 c512T→A SMEI Missense T199R c596C-→G 4, 29 SMEI⁴ Missense c677C→T T226M 5,30 SMEI Missense c715G→A A239T 6,31 SMEI Missense c2837G→A R946H 7.32 SMET Missense c3714A→C E1238D 8.33 SMEI Missense c4186T→G C1396G 9, 34 SMEI Missense c4321G→C A1441P 10.35 SMET Missense 11545V 11,36 c4633A→G SMEI Missense c4934G→A R1645Q 12, 37 SMEI Missense c5119T→G F1707V 13.38 SMEI Missense A1783T c5347G→A 14, 39 SMET Truncation c41delT F14fsX91 15.40 SMET Truncation c496insGTGAATC T166fsX170 16, 41 SMEI Truncation c1687delC L563fsX622 17, 42 SMEI Truncation c3231delA K1077fsX1079 18, 43 SMEI Truncation c3561-3562delAA O1187fsX1215 19,44 SMEI Truncation c4062delT C1354fsX1359 20, 45 N1509fsX1511 SMEI Truncation c4526delA 21,46 SMEI Nonsense c3022G→T E1008X 22, 47 SMEI Nonsense c4279C→T Q1427X 23, 48 SMEI3 Splice Site IVS4+5G→A 24 SMEI3 Splice Site 25 IVS3-13T→A Non-SMEI Missense c580G→A D194N 49. 54 Missense Non-SMEI c4439G→T G1480V 50, 55 Non-SMEI Missense c4907G→A R1636O 51, 56 Non-SMEI Truncation c1724delT F575fsX622 52, 57 Non-SMEI Truncation c5741-5742delAA O1914fsX1943 53, 58

Note: "Patient diagnosis was based on the initial clinical observations. "Numbering is based on the large SCN1A isoform." These splice site alterations were seen in the same individual. "This atteration was also seen in an individual that was not clinically diagnosed with SMEI. WO 2004/085674 PCT/AU2004/000295 - 45 -

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PCT/AU2004/000295

Claims

1. A method for the diagnosis of SMEI in a patient comprising detecting an alteration in the SCM1A gene, including in a regulatory region of the gene, in a patient sample, and ascertaining whether the alteration is known to be SMEI associated or non-SMEI associated or, if not known to be either, determining the likelihood that it is a SMEI associated alteration.

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 A method as claimed in claim 1 further comprising establishing a diagnosis which will indicate a high probability of SMEI where the alteration is known to be SMEI associated.

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- 3. A method as claimed in claim 2 wherein the alteration is one of those alterations identified as SMEI associated in Table 3.
- 20 4. A method as claimed in claim 1 comprising establishing a diagnosis which will indicate a low probability of SMEI where the alteration is non-SMEI associated.
- 25 5. A method as claimed in claim 4 wherein the alteration is one identified of those alterations identified as non-SMEI associated in Table 3.
- 6. A method as claimed in claim 1 wherein the likelihood that the alteration is a SMEI associated alteration is established through:

considering genetic data for parents and/or relatives; and

establishing whether the alteration has arisen de 35 novo or is inherited.

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- 7. A method as claimed in claim 6 further comprising establishing whether the alteration would result in a major disruption to the protein.
- 5 8. A method as claimed in claim 7 wherein the alteration is a truncating mutation.
- 9. A method as claimed in any one of claims 6 to 8 comprising establishing a diagnosis which will indicate a low probability of SMEI in the case of an inherited mutation and indicate a high probability of SMEI in the case of a de novo mutation, and a very high probability of SMEI where a de novo mutation would result in a major disruption to the protein.
 - 10. A method as claimed in any one of claims 1 to 9 comprising performing one or more assays to test for the existence of an SCNIA alteration and to identify the nature of the alteration.
 - 11. A method as claimed in claim 10 comprising:

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- performing one or more assays to test for the existence of an alteration in the SCN1A gene of the patient; and, if the results indicate the existence of an alteration in the SCN1A gene,
- (2) performing one or more assays to identify the nature of the SCNIA alteration.
- 12. A method as claimed in claim 10 or 11 wherein one of 30 the assays is a DNA hybridisation assay.
- 13. A method as claimed in claim 12 wherein an SCN1A gene probe, an SCN1A exon-specific probe, or an SCN1A allele specific probe is hybridised to genomic DNA isolated from standard patient.

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allele specific oligonucleotides.

14. A method as claimed in claim 10 or 11 wherein one of the assays is high performance liquid chromatography.

- 15. A method as claimed in claim 10 or 11 wherein one of the assays is an electrophoretic assay.
- 16. A method as claimed in claim 10 or 11 wherein the sample DNA to be tested is quantitatively amplified for at least one exon of the SCNIA gene to produce amplified fragments and the length of the amplification products for each amplified exon is compared to the length of the amplification products obtained when a wild-type SCNIA gene is amplified using the same primers, whereby differences in length between an amplified sample exon and the corresponding amplified wild-type exon reflect the
- 17. A method as claimed in claim 10 or 11 wherein one of the assays incorporates DNA amplification using SCNIA

occurrence of a truncating alteration in the sample SCN1A

- 19. A method as claimed in claim 10 or 11 wherein one of the assays is SSCP analysis.
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gene.

- 19. A method as claimed in claim 10 or 11 wherein one of the assays is RNase protection.
- 20. A method as claimed in claim 10 or 11 wherein one of 30 the assays is DGGE.
 - 21. A method as claimed in claim 10 or 11 wherein one of the assays is an enzymatic assay.
- 35 22. A method as claimed in claim 21 wherein said assay incorporates the use of MutS.

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23. A method as claimed in claim 10 or 11 wherein one of the assays examines the electrophoretic mobility of the SCNIA protein of the patient.

- 5 24. A method as claimed in claim 10 or 11 wherein one of the assays is an immunoassay.
 - 25. A method as claimed in claim 10 or 11 wherein one of the assays is DNA sequencing.

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26. A method for the diagnosis of SMEI in a patient, comprising:

detecting an alteration in the SCNIA gene, including in a regulatory region of the gene, in a patient sample; 15 and

establishing a diagnosis which will indicate a high probability of SMEI if a SMEI associated alteration as laid out in Table 3 is identified or, in the alternative, establishing a diagnosis which will indicate a low

- 20 probability of SMEI if a non-SMEI associated alteration as laid out in Table 3 is identified.
 - 27. A method of determining the appropriate treatment for a SMEI patient comprising performing one or more of the methods claimed in any one claims 1 to 26 and correlating the diagnosis reached with known indications and contraindications for SMEI patients.
- 28. A method of determining the likelihood of adverse results from treatments of a SMEI patient including drug treatments and vaccinations comprising performing one or more of the methods as claimed in any one of claims 1 to 26 and correlating the diagnosis reached with known indications and contra-indications for SMEI patients.
 - 29. An isolated nucleic acid molecule encoding an altered SCN1A subunit of a mammalian voltage-gated sodium channel,

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wherein the alteration gives rise to an SMEI phenotype and has the sequence set forth in any one of SEQ ID NOS: 1-25.

- 30. An isolated nucleic acid molecule encoding an altered 5 SCNIA subunit of a mammalian voltage-gated sodium channel, wherein the alteration gives rise to a non-SMEI epilepsy phenotype and has the sequence set forth in one of SEQ ID NOS: 49-53.
- 31. An isolated nucleic acid molecule comprising the nucleotide sequence set forth in any one of SEQ ID NOs: 1-25, 49-53.
- 32. An isolated nucleic acid molecule consisting of the nucleotide sequence set forth in any one of SEQ ID NOs: 1-25, 49-53.
 - 33. An expression vector comprising a nucleic acid molecule as claimed in any one of claims 29 to 32.
- 34. A cell comprising a nucleic acid molecule as claimed in any one of claims 29 to 32.

- 35. A genetically modified non-human animal comprising a 25 nucleic acid molecule as claimed in any one of claims 29 to 32.
- 36. A genetically modified non-human animal as claimed in claim 35 in which the animal is selected from the group 30 consisting of rats, mice, hamsters, guinea pigs, rabbits, dogs, cats, goats, sheep, pigs and non-human primates such as monkeys and chimpanzees.
 - 37. An isolated polypeptide, said polypeptide being an altered SCN1A subunit of a mammalian voltage-gated sodium channel, wherein the polypeptide has the amino acid

sequence set forth in one of SEQ ID NOS: 26-48 and the alteration gives rise to an SMEI phenotype.

- 38. An isolated polypeptide, said polypeptide being an altered SCMIA subunit of a mammalian voltage-gated sodium channel, wherein the polypeptide has the amino acid sequence set forth in one of SEQ ID NOS: 54-58 and the alteration gives rise to a non-SMEI epilepsy phenotype.
- 39. An isolated polypeptide comprising the amino acid sequence set forth in any one of SEQ ID NOs: 26-48, 54-58.
 - 40. An isolated polypeptide consisting of the amino acid sequence set forth in any one of SEQ ID NOs: 26-48, 54-58.
 - 41. A sodium channel that incorporates an SCNIA subunit as claimed in any one of claims 37 to 40.
- 42. A method of preparing a polypeptide comprising the 20 steps of:
 - 1) culturing a cell as claimed in claim 34 under conditions effective for polypeptide production; and
 - 2) harvesting the polypeptide.

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- 25 43. A polypeptide prepared by the method of claim 42.
 - 44. An antibody which is immunologically reactive with an altered polypeptide as claimed in any one of claims 37 to 40 or 43, or a sodium channel as claimed in claim 41, but not with a wild-type sodium channel.
 - 45. An antibody as claimed in claim 44 which is selected from the group consisting of a monoclonal antibody, a humanised antibody, a chimeric antibody or an antibody fragment including a Fab fragment, (Fab')2 fragment, Fv fragment, single chain antibodies and single domain antibodies.

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46. The use of a nucleic acid molecule as claimed in any one of claims 29 to 32 for the screening of candidate pharmaceutical compounds.

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47. The use of a polypeptide as claimed in any one of claims 37 to 40 or 43, a sodium channel as claimed in claim 41, or an antibody as claimed in claim 44 or 45 for the screening of candidate pharmaceutical compounds.

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- 48. The use of genetically modified non-human animal as claimed in claim 35 or 36 or a cell as claimed in claim 34 in the screening of candidate pharmaceutical compounds.
- 49. A compound when identified through a use as claimed in any one of claims 46 to 48.
 - 50. A pharmaceutical composition comprising a compound as claimed in claim 49 and a pharmaceutically acceptable carrier.
 - 51. A method of treating epilepsy, including SMEI, comprising administering a selective antagonist, agonist or modulator of a polypeptide as claimed in any one of claims 37 to 40, or a sodium channel as claimed in claim 41 to a patient in need of such treatment.
- 52. The use of a selective antagonist, agonist or modulator of a polypeptide as claimed in any one of claims 30 37 to 40, or a sodium channel as claimed in claim 41 in the manufacture of a medicament for the treatment of epilepsy, including SMEI.
 - 53. A method of treating epilepsy, including SMEI, comprising administering an isolated nucleic acid molecule which is the complement (antisense) of a nucleic acid molecule as claimed in any one of claims 29 to 32 and

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which encodes an RNA molecule that hybridizes with the mRNA encoding an altered SCNIA protein to a subject in need of such treatment.

- 5 54. The use of an isolated nucleic acid molecule which is the complement (antisense) of a nucleic acid molecule as claimed in any one of claims 29 to 32 and which encodes an RNA molecule that hybridizes with the mRNA encoding an altered SCNIA polypeptide in the manufacture of a medicament for the treatment of epilepsy, including SMBI.
 - 55. A method of treating epilepsy, including SMEI, comprising administration of an antibody as claimed in claim 44 or 45.
 - 56. The use of an antibody as claimed in claim 44 or 45 in the manufacture of a medicament for the treatment of epilepsy, including SMEI.

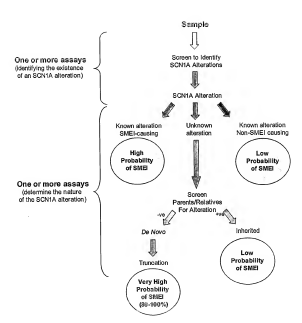
- 57. A method of treating epilepsy, including SMEI, comprising administering an antibody, as claimed in claim 44 or 45, administration of an agonist, antagonist or modulator of a polypeptide as claimed in any one of claims 37 to 40, or a sodium channel as claimed in claim 41, or 25 administration of a DNA molecule which is the complement of a nucleic acid molecule as claimed in any one of claims 29 to 32 and which encodes an RNA molecule that hybridizes with the mRNA encoding an altered SCNIA protein, in combination with administration of the wild-type SCNIA, to a subject in need of such treatment.
 - 58. The use of an antibody, as claimed in claim 44 or 45, use of an agonist, antagonist or modulator of a polypeptide as claimed in any one of claims 37 to 40, or a sodium channel as claimed in claim 41, or use of a DNA molecule which is the complement of a nucleic acid molecule as claimed in any one of claims 29 to 32 and

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which encodes an RNA molecule that hybridizes with the mRNA encoding an altered SCNIA protein, in combination with the use of the wild-type SCNIA, in the manufacture of a medicament for the treatment of epilepsy, including SMEI.

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Figure 1



INTERNATIONAL SEARCH REPORT

International application No.
PCT/AU2004/000295

A.	CLASSIFICATION OF SUBJECT MATTER					
Int. C1. 7:	C12Q 1/68, C12N 15/01, A61K 39/395, C	7K 14/47				
According to International Patent Classification (IPC) or to both national classification and IPC						
B.	B. FIELDS SEARCHED					
	Minimum documentation searched (classification system followed by classification symbols) SEE ELECTRONIC DATABASES					
	Documentation searched other than minimum documentstion to the extent that such documents are included in the fields searched SEE ELECTRONIC DATABASES					
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WIPS, Medline, CA (see continuation of box B)						
C.	DOCUMENTS CONSIDERED TO BE RELEVAN					
Category ¹	Citation of document, with indication, where	ppropriate, of the relevant passages	. Relevant to claim No.			
X	Fujiwara, T. et al. 2003. Mutations of se intractable childhood epilepsies with free a Journal of Neurology, Vol: 126, pages	uent generalised tonic-clonic scizures. I				
х	Lerche, H. et al. 2001. Ion channels and epilepsy. American Journal of Medical Genetics, Vol: 106, pages 146-159.					
X .	Cannon, Stephen.C. 2002. Sodium char. 34, pages 853-858.	nel gating: no margin for error. Neuron,	Vol: 1-2, 4 and 10-11			
X	Further documents are listed in the continua	ion of Box C See patent fam	ily annex			
"A" docu	ial categories of cited documents: ment defining the general state of the art which is "T" onsidered to be of particular relevance	later document published after the international filing conflict with the application but cited to understand a underlying the invention				
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or w	ment which may throw doubts on priority claim(s) "Y" nich is cited to establish the publication date of	alone document of particular relevance; the claimed inventi involve an inventive step when the document is comb	ined with one or more other			
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C (Continuat	ion). DOCUMENTS CONSIDERED TO BE RELEVANT	
Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	Sugawara, T. et al. 2002. Frequent mutations of SCN1A in severe myclonic epilepsey in infancy. Neurology, Vol. 58, pages 1122-1124.	1-2, 4, 6-25 and 27-28.
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х	Malacame, M. et al. 2002. Lack of SCN1A mutation in familial febrile seizures. Epilepsia, Vol: 43(5), pages 559-562.	1-2, 4 and 6-25
х	Ohmori, I. et al. 2002. Significant correlation of the ScN1A mutations and severe myclonic epilepsy in infancy. Biochemical and Biophysical Research Communications, Vol. 295, pages 17-23.	1-2, 4, 6-25
A	Spampanato, J. et al. 2003. Generalised epilepsy with febrile seizures plus type 2 mutation W1204R alters voltage-dependent gating of Na,1.1 sodium channels. Neuroscience, Vol: 116, pages 37-48.	1-2 and 4

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Supplemental Box (To be used when the space in any of Boxes I to VIII is not sufficient)
Continuation of Box No: B (Search Terms)
Epilepsy, Sodium channel, SCN1A, Severe myclonic epilepsy of infancy, SMEI, mutation, allele,
Polymorphism, antibody, anti-sense, ribosyme, RPAi, agonist, antagonist, modulate, inhibit.